

## **Predictive Factors of Clinical Assays during COVID-19**

Matthieu MILLION, Pierre DUDOUE, Eric CHABRIERE, Sébastien CORTAREDONA,  
Philippe BROUQUI, Didier RAOULT

### **SUPPLEMENTARY DATA**

#### **SUPPLEMENTARY METHODS**

##### *Global strategy*

In the context of a global crisis of trust in medical literature triggered by a retracted article published in the Lancet (1), we performed a critical reading of scientific publications on the clinical efficacy of chloroquine derivatives and remdesivir against Covid-19 since March 2020 (2). We primarily focused on mortality and considered all studies with Covid-19 patients treated or not treated by hydroxychloroquine (HCQ) or remdesivir with at least 1 death. We began by looking at each article and identifying anomalies that we felt were unacceptable or to be avoided from a medical point of view (3-7). Gradually this led us to identify essential or recommended judgement criteria which were gathered in a checklist. In November 2020, we stopped this checklist, which is provided here (Table 1). We then comprehensively reviewed all the publications and preprints with this checklist and described for each criterion the triggering study, and all the studies that did not meet them. We then analyzed all the articles using unsupervised approach. Finally, we performed a comparative meta-analysis, as described previously (2), comparing studies that met or did not meet each criterion. When 2 studies studied common patients or the same cohort, both of them could be analyzed for criteria identification but only the most recent one, with the largest number of patients, published versus preprint or including the most recommended criteria identified here were included in the quantitative meta-analysis to assess HCQ efficacy.

### ***Inclusions of studies: Search strategy***

The keywords “hydroxychloroquine”, “HCQ”, “chloroquine”, “coronavirus”, “COVID-19” and “SARS-Cov-2”, “remdesivir” were used in the PubMed, Google Scholar and Google search engines for studies published in English (research updated on November, 11, 2020). An online search was also performed using the website <https://c19study.com/>. The following outcome was considered: death, so studies without any death were not eligible. Preprints were also included. When preprints were subsequently published, final publication and preprints were compared. We reviewed studies evaluating the effects of chloroquine derivatives against SARS-CoV-2 in groups of COVID-19 patients compared to control groups of patients who did not receive chloroquine derivatives. Articles published in peer-reviewed journals, preprints and articles available on the internet, even when not published on official websites, were included. Manuscripts submitted to a peer-reviewed journal but not published online and whose submitted draft leaked on the internet were not included. Only studies comparing a group of COVID-19 patients treated with a chloroquine derivative to a control group without chloroquine derivatives were included. Noncomparative (single arm) studies and studies comparing two groups treated with chloroquine derivatives at different dosages or with different delays of treatment were not eligible. Studies analyzing safety, efficacy as a prevention, and data provided as a webpage without any article format (such as a tweet), were also not eligible.

### ***Identification of characteristics and criteria***

The criteria are summarized in Table 1. Some of these criteria have already been identified in a previous work (3,8) and have been completed as we observed critical pitfalls in studies assessed for the present work. A criterion was not fulfilled if it was mentioned but not fulfilled and/or if it was not mentioned.

In the retracted article (1) which triggered the scandal, we identified several quality criteria not fulfilled: *Absence of private company computing data, Centers and doctors who take care of patients are identified, The therapeutic protocol is detailed (standard care, evaluation of contraindications, dosage and duration)* and *At least one main author is a clinical expert-in-the-field (affiliated to an infectious disease, internal medicine or a pneumology unit)*. Indeed, a private data computing company collected data (Surgisphere), centers and doctors were not identified, therapeutic protocol was not mentioned, and authors were affiliated to biomedical or heart and vascular units.

In other studies, we identified the following medical quality criteria: *Potential conflict of interest* such as a study reporting an increased mortality with HCQ compared with standard-of-care funded by the company marketing remdesivir (9), with a design strikingly similar to the retracted article (1). Potential conflict of interest was defined when the name of a company marketing remdesivir was mentioned in the manuscript as a funder or as a conflict of interest with at least 1 author or 1 investigator either declared or found on transparency websites (transparence.sante.gouv.fr, eurosfordocs, dollarsfordocs) but not declared. A non-compensated consulting was not considered a potential conflict of interest (10). *Absence of undeclared funding and conflict of interest*: an author disclosed a financial relationship with a company marketing remdesivir in 2019 (<https://www.astmh.org/ASTMH/media/2019-Annual-Meeting/ASTMH-2019-Speaker-Disclosure-Statement.pdf>), but not in his two studies reporting an absence of effect of HCQ to prevent (11) or treat Covid-19 (12). *Patients without confirmation of diagnosis by a microbiological test are excluded*: The same authors not declaring any conflict of interest confirmed cases with a microbiological test in only 18% (11) or 34% of cases (12). Laboratory confirmation is essential as clinical diagnosis is not sufficient as many respiratory viruses circulate at the same time (13). *The treatment is not toxic (not overdosed or used in contraindicated patients)*: A study (14) used 1.5 times the

loading dose of chloroquine-sensitive malaria ([www.cdc.gov](http://www.cdc.gov)), and 4 times the usual dosage in other acute infectious diseases, such as liver amebiasis (15). ***Patients in the no-treatment group are not treated with the experimental treatment or with any other treatment that the treated group did not have.*** This was observed in a study in which treated patients received only HCQ but 30% of untreated patients received azithromycin (16). ***Confounding role of previous health status (at least age) is ruled out.*** This was not the case in a paper (17) where patients were older, but no attempt was done to control this confounding. ***Confounding role of disease severity (at least oxygen status) is ruled out.*** Strikingly, in a study (9), twice as many patients were intubated in the HCQ group than in the non-HCQ group (24.9% vs 12.2%) and this was not controlled. We already commented this (5). Other mistakes were observed but their effect could not be adequately quantified by quantitative meta-analysis and Q-value. ***Conclusions neglecting a non-significant decrease or increase in mortality of 25% or more*** (18). In this case, indeed, there is a difference but as the study does not have the power to confirm it significantly, it could be due to chance or to the poor design. Typically, these studies should be used for meta-analysis that will confer the power to confirm or not the significance of the difference. ***Conclusions neglecting an unexpected relevant result*** : A study found no death in patients treated by the combination therapy associating hydroxychloroquine and azithromycin (19), but this was not tested nor discussed. ***The main outcome is objective, independent of human subjectivity and context and did not change during study:*** In an observational study, the main outcome was death and/or transfer to intensive care unit (ICU) (20) but ICU transfer is highly subjective and depends on the physician and the number of available ICU beds. Death is only mentioned in the supplementary data without methods to control confounding with previous health status or severity while treated patients were much more severe at baseline. In an RCT (21), the main outcome changed from “difference in clinical status” to “time to recovery” during the study.

***Identified articles: preprints, published articles, censorship during editing***

Overall, 61 studies were evaluated. For HCQ/CQ, 56 studies (with at least 1 death) were identified (Supplementary File 1) corresponding to 23 preprints (14 without publication in a (peer-reviewed) journal, 9 preprints subsequently published in a journal), and 33 studies published in a journal without previous preprint. A preprint (22) and a study published in a journal (23) from different authors analyzed the same Spanish cohort. For remdesivir, only 6 studies were found including 3 preprints and 4 peer-reviewed publications (10, 21,24-28). One study was common for HCQ and remdesivir and was published both as a preprint (24) and a peer-reviewed publication (25).

We observed discordances between preprints and final manuscripts. Data evidencing a favorable effect of HCQ (alleviations of symptoms, greater reduction of CRP, more rapid recovery from lymphopenia) were mentioned in the preprint (29) but removed in the final published version (30). This deletion was requested by the editor of the journal. Conversely, Magagnoli improved quality between preprint (16) and final publication (31) including a subgroup analysis by severity before treatment.

The 56 studies on HCQ/CQ came from the USA (16 studies), France (n = 9), Spain (n = 6), Italy (n = 4), Iran, Ireland (2 studies each), and Andorra, Belgium, Brazil, China, Congo, Egypt, Greece, India, Mexico, the Netherlands, Peru, Saudi Arabia, Turkey and the United Kingdom (1 each). Three involved more than 1 country. Strikingly, only 1 included study came from China while several comparative studies have been reported from this country without any death. For remdesivir, 3 studies were performed in the USA, 1 in China, 1 in Poland and 1 was multinational.

Among all 61 evaluated studies, 49 were observational including 24 Big data studies. We found 12 RCTs including 9 megatrials. Forty-three studies were multicentric and 18

were monocentric. For 6 studies, data for death analysis were not sufficient for quantitative meta-analysis (sample size in each group, with number of death or summary result for death not provided).

### **Funding, conflict of interest of studies evaluating HCQ or remdesivir on Covid-19 mortality**

We considered it to be a conflict of interest when the study was funded by Gilead directly (remdesivir) or indirectly (9) or when at least 1 author received fees from Gilead and declared it or did not declare it.

#### *Studies funded by pharmaceutical industries*

We found that 4 studies were funded by pharmaceutical industries. Studies by Fried et al. (9), Goldman et al. (32) and Spinner et al. (28) were funded by Gilead who market remdesivir. Cavalcanti (33) was funded by the first Brazilian big pharma industrial (EMS Pharma) but we found no link about this industrial regarding a conflict of interest so this study was considered “without conflict of interest”. These 4 studies were published in the journals with the highest impact factors in medicine and infectious diseases. In the RCT reported by Goldman (32) comparing two durations of remdesivir (without placebo), 109/397 (27.4%) patients were treated with HCQ and mentioned in supplementary data but were not analyzed. In this RCT, hydroxychloroquine was associated with lower death rate (9 versus 12%).

#### *Declared conflict of interests*

In Biegel et al. (21), employees of Gilead Sciences participated in discussions about protocol development and in weekly protocol team calls. Seven authors declared a conflict of interests

with Gilead. In Flisiak et al. (27), 6 / 22 authors received personal fees from Gilead and this was declared.

#### *Undeclared conflict of interests*

In Geleris et al. (20), an author received at least 9,413\$ for consulting from Gilead Sciences inc on Jan 31, 2018 (<https://projects.propublica.org/docdollars/>).

Mahevas et al. (34) declared no funding received nor conflict of interest in their study but the competing interests were not fully declared in the original publication so that an erratum was published (35) with an updated and expanded conflict of interest statement with almost all authors receiving personal fees from pharmaceutical industries. Another author of the same work, with initially undeclared conflict of interest in this publication, declared a conflict of interest in some publications on HCQ and remdesivir (36) but not in others (37-40).

An author (D. Boulware) disclosed a financial relationship with Gilead in 2019 (<https://www.astmh.org/ASTMH/media/2019-Annual-Meeting/ASTMH-2019-Speaker-Disclosure-Statement.pdf>), but not in his two studies reporting an absence of effect of hydroxychloroquine to prevent (11) or treat Covid-19 (12).

In the WHO Solidarity trial (24,25) published as a preprint in MedRxiv, no author declared any conflict of interest while in supplementary data, it appeared that several participants, especially investigators who included patients in the trials, had received fees from Gilead. Moreover, 4 authors finally reported personal fees from Gilead in the final publication (25) whereas this was not reported in the preprint where it could be read “Competing Interest Statement: The authors have declared no competing interest’ (24).

#### *Possible conflict of interests*

In two articles, we found several conflicts of interests between authors and several pharmaceutical industries (41,42), however, we did not find Gilead in these industries. It is however, possible that unreported conflicts of interests exist between other firms and a possible efficacy of hydroxychloroquine.

Besides for-profit private data computing companies, we found two Big data studies performed with the US Department of veterans affairs associated with HCQ inefficacy (26,43-44) and remdesivir efficacy (26). Strikingly, Gilead supports veterans through the Gilead Veterans Engagement Team (<https://www.gilead.com/careers/inclusion-and-diversity>) and has intricated relationships with the US Department of veterans affairs since anti-HCV sofosbuvir development (<https://www.military.com/daily-news/2016/02/05/former-va-scientist-responds-to-lawmakers-suspicious-drug-sale.html>). Furthermore, Gilead provided remdesivir to US army at no cost (<https://www.militarytimes.com/news/your-military/2020/03/10/army-signs-agreement-with-drug-giant-gilead-on-experimental-covid-19-treatment/>).

### **For-profit private data computing companies and big data studies**

We found 3 big data studies with a possible shell company (private data computing company). Surgisphere was a private data computing company in a study subsequently retracted (1). We did not succeed to identify main actionnaires of this company despite thorough internet research (<https://www.prnewswire.com/news-releases/ihfs-global-healthcare-quality-award-recognizes-surgisphere-executive-sapan-desai-md-300637851.html>). TARGET PharmaSolutions in a study published in Clinical Infectious Diseases with funding for initial data acquisition provided by Gilead (9), and TriNetX in a preprint (43) and in a published paper (44). Target PharmaSolutions is a for-profit company with a total funding amount of \$637K with 5 members and 3 investors funded by the first author of the publication (M. Fried



(<https://www.crunchbase.com/organization/target-pharmasolutions>), (9)). TriNetX is an initiative of the West Virginia Clinical and Translational Science Institute (<https://www.wvctsi.org/programs/epidemiology-biostatistics/trinetx/>), with active link with Sanofi (<https://trinetx.com/sanofi/>), Merck, Itochu, Novartis, and Pfizer (<https://www.outsourcing-pharma.com/Article/2018/01/16/Sanofi-partners-with-TriNetX-to-speed-drug-development-timelines> & <https://www.frenchweb.fr/trinetx-leve-40-millions-de-dollars-pour-exporter-ses-solutions-doptimisation-des-essais-cliniques-en-europe/351399>).

### **Studies that did not mention treatment details**

Contraindications are not mentioned in several big data studies (22). In the big data study by Sbidian et al. (45) including 39 hospitals in Paris, it is not possible to know the posology nor the duration. The suggested HCQ regimen is mentioned “loading dose of 600 mg on day 1, followed by 400 mg daily for 9 additional days. AZI at a dose of 500 mg on day 1 and then 250 mg daily for 4 more days in combination with HCQ was an additional suggested therapeutic option. Prescription of HCQ or HCQ together with AZI was at the discretion of the physicians.” In this multicentric big data study, the absence of data on treatment and the absence of standardized protocol may prevent any conclusion.

### **Studies without control for initial disease severity**

#### *Eight studies with treated patients more severe at baseline*

We found 8 studies in which severity was not controlled for and with treated patients more severe than untreated patients. Strikingly, in the study by Fried et al. (9), whose initial data acquisition was provided by Gilead, HCQ-treated patients were more severe (more frequent pneumonia) and the authors reported an increased mortality in the HCQ group without adjusting for any confounding. In Geleris *et al.* (20) published in the NEJM, the use of

propensity score was not sufficient and after matching the treated group still had a 20% lower PaO<sub>2</sub>/FiO<sub>2</sub> ratio, a 40% higher ferritin, and an 18% higher CRP than the untreated group. In the study by Ip *et al.* (41) HCQ-treated patients were almost 2 times more likely to have a SaO<sub>2</sub> < 94% (49% vs 30%, p < 0.05), and the propensity score model did not include this parameter while age, comorbidities and “log ferritin” were included in the model. In Kelly *et al.* (46) HCQ-treated patients had significantly higher CRP, FiO<sub>2</sub> requirement and clinical scale at day 0 and there was no attempt to control this confounding. Magagnoli *et al.* (16,31) reported a propensity score analysis without mentioning covariates included in the model. Because treated patients were much more severe (lymphopenia twice as common in the treated group (25%) than in the untreated group (14%)), it was not possible to rule out a confounding role of severity. McGrail (17) reported that treated patients were older and more severe but did not attempt to control these confoundings. Finally, in the study by Peters *et al.* (47) treatment was started when there was an increase in respiratory rate or use of supplemental oxygen. This implied an uncontrollable confounding by indication bias. This “confounding by indication” bias seems associated with big data as we also found that severity was not adequately adjusted for in the study of the Covid-19 cancer consortium (matched data presented in Supplemental Table S5 of Rivera *et al.* (42): 93% moderate-severe in the HCQ group versus 80% in the untreated group).

A conflict of interest was found for 3 of these studies (9,16,20,31) and highly suspected for 2 of them (41,42).

*We did not find any study in which the treated patients were less severe than the untreated ones.*

*Studies in which difference of severity between treated and untreated could not be assessed*

We found 16 studies in which a difference in severity between treated and untreated was not evidenced but could not be ruled out. Alamdari *et al.* reported that expired patients presented more frequently with shortness of breath at admission and were less frequently treated, however effect of treatment was not controlled for initial severity (48). In Alberici *et al.* (18) HCQ was associated with an important protective effect against death (OR = 0.44,  $p > 0.05$ ) but HCQ was not included in multivariate analyses because p-value was not  $< 0.5$ . Indeed, only the statistically-significant predictors at univariate analysis were entered into a multivariate model. Bhandari *et al.* (49) reported, among asymptomatic patients at inclusion, 1 death /39 in the HCQ group versus 1/32 in the control one, however, oximetry was not provided in any of the two groups. As hypoxia could be asymptomatic (50), a difference in initial severity could not be ruled out. In the same study (50), asymptomatics were treated with HCQ or no treatment, mild ill were treated with HCQ, severely ill with HCQ AZ and critically ill with Lopinavir+ritonavir so that it was not possible to control for the role of disease severity. Calik Basaran *et al.* (51) reported a shorter length of hospitalization in HCQ AZ but severity between groups was different at baseline and exposition of the 4 dead people (treated or untreated) was not provided. Derwand *et al.* provided no information on the control population (52). Heberto *et al.* (53) reported a significantly decreased mortality in multivariate analysis but potential predictors included in the model were not provided, notably because myocardial injury but not death was the main outcome. Goldman (32) found a mortality decrease with HCQ but did not analyze it because it was not the main outcome as the study was designed to assess remdesivir. Guerin *et al.* (54) performed a case-control sub-analysis matched for age, sex, and body mass index but not severity while some patients were severe (respiratory rate ranging from 12 to 50). Some studies reporting multivariate analyses did not mention the covariates included in the models, so a role of severity could not be

excluded (54). Pinato *et al.* made no mention of disease severity or oxygen requirements (55). In Roomi *et al.* (56), age was not different and controlled for in multivariate analyses. However, initial disease severity was not assessed and not included in multivariate analysis. Serrano in their abstract did not mention baseline characteristics and did not attempt to control for age or severity (57). In Singh *et al.* previous health status and comorbidities were included for matching but disease severity was not considered in the propensity matching (43). Skipper *et al.* in their internet-based RCT assessed “shortness of breath” but did not assess oxygen status (oximetry) at baseline (12). Soto-Beccera *et al.* developed a complex model including several comorbidities and “pneumonia diagnosed within 48 hours of admission” but not oxygen status (58). Because we treated more than 10,000 patients in our center, it is clear that pneumonia could be minimal, intermediate or severe with a very different risk of complications between minimal (<10% lung volume) and severe (>50%) involvement (59,60). Furthermore, since hypoxia is frequently asymptomatic (51), oxygen status could not rely on interview but required objective measurement. Sulaiman performed multivariate analysis including age, gender and comorbidities but disease severity was not controlled for (61). Synolaki did not analyze confounding for treatment as it was not the main topic of the paper (62).

**Supplementary Table 1. Comparative meta-analysis according to quality criteria identified in the present study**

	Q-value	p-value of Q-value	Summary effect of studies fulfilling this criterion				Summary effect of studies not fulfilling this criterion			
			number of comparisons	OR	95%CI	p-value	number of comparisons	OR	95%CI	p-value
<b>Absence of private data computing company</b>	73.1	<.0001	56	0.81	0.74 - 0.89	<.0001	6	1.28	1.23 - 0.34	< .0001
<b>Potential conflict of interest</b>	39.3	< .0001	43	0.75	0.66 - 0.83	< .0001	19	1.15	1.07 - 1.23	0.0001
<b>Detailed therapeutic protocol</b>	28.5	< .0001	25	0.68	0.59 – 0.78	< .0001	37	1.04	0.96 – 1.12	0.34
<b>Centers and doctors who take care of patients are identified</b>	27.2	< .0001	41	0.70	0.61 - 0.80	< .0001	21	1.09	0.99 - 1.21	0.07
<b>An author clinical expert-in-the-field*</b>	21.0	< .0001	29	0.71	0.61 – 0.81	< .0001	25	1.06	0.96 – 1.16	0.25
<b>Clinical study (Not a 'Big data' study based on electronic medical files)</b>	14.5	0.0001	34	0.66	0.55 – 0.79	< .0001	28	0.99	0.89 – 1.09	0.83
<b>Nontoxic treatment (dose, use in contraindicated patients)</b>	13.9	0.0002	58	0.85	0.78 - 0.93	0.0008	4	1.09	0.998 - 1.20	0.06

<b>Observational (versus interventional) (RCTs)</b>	12.5	0.0004	52	0.85	0.77 – 0.94	0.001	10	1.07	0.98 – 1.17	0.11
<b>Monocentric (versus multicentric)</b>	12.3	0.0004	18	0.55	0.41 – 0.73	< .0001	44	0.95	0.87 – 1.03	0.22
<b>Not a megatrial</b>	11.9	0.001	55	0.86	0.78 – 0.94	0.001	7	1.07	0.98 – 1.17	0.11
<b>Control group without another specific treatment effective on SARS-Cov-2 (other treatment, HCQ or AZ)</b>	11.4	0.001	60	0.85	0.78 - 0.93	0.001	2	1.33	1.05 - 1.70	0.02
<b>Number of events, total treated untreated known</b>	7.66	0.006	49	0.94	0.86 - 1.03	0.18	13	0.67	0.54 - 0.83	0.0004
<b>Treatment monitoring</b>	7.43	0.006	19	0.70	0.59 – 0.84	0.0001	43	0.93	0.85 – 1.03	0.16
<b>Funding is mentioned, absence undeclared COI</b>	7.11	0.008	55	0.86	0.78 - 0.94	0.001	7	1.07	0.93 - 1.23	0.32
<b>Control for severity (at least oxygen)</b>	6.62	0.01	39	0.80	0.72 - 0.90	0.0001	23	1.02	0.88 - 1.18	0.79
<b>Absence of mixed stages of the disease</b>	6.52	0.01	39	0.79	0.70 – 0.89	0.0001	23	0.98	0.87 – 1.10	0.72
<b>Detailed Standard of care (Soc)</b>	6.03	0.01	7	0.60	0.45 – 0.82	0.001	55	0.90	0.82 – 0.986	0.023
<b>Diagnosis formally confirmed (PCR or serology-based diagnosis)</b>	4.98	0.026	48	0.84	0.76 - 0.93	0.001	14	1.04	0.89 - 1.21	0.64

<b>Conclusions do not neglect a 25% difference in mortality risk</b>	1.51	0.22	51	0.87	0.79 – 0.96	0.004	11	0.98	0.83 – 1.14	0.76
<b>Unexpected findings reported</b>	0.86	0.35	56	0.86	0.78 – 0.94	0.001	6	0.998	0.74 – 1.35	0.991
<b>Objective outcome</b>	0.23	0.63	56	0.87	0.79 – 0.95	0.002	6	0.90	0.80 – 1.002	0.053
<b>Control for health status (at least age)</b>	0.047	0.83	55	0.87	0.79 - 0.95	0.002	7	0.82	0.48 - 1.39	0.45

Random effect model, 62 comparisons. \*For 8 comparisons, this could not be determined.

**Supplementary Table 2. Criteria identified through errors and mistakes in analysis of studies assessing HCQ and remdesivir for Covid-19**

Criteria	Explanation	PRISMA Checklist	STROBE Checklist	CONSORT Checklist
<b>Potential sources of bias</b>	<p>In usual checklists, potential sources of bias are mentioned but not identified.</p> <p>In the context of Covid-19, the major sources of biases identified in the present study were conflict of interest and lack of clinical expertise.</p>	No	<p>Item 9. Describe any efforts to address potential sources of bias</p> <p><i>These sources of bias are not clearly identified. Conflict of interest and lack of clinical expertise not considered as potential sources of bias.</i></p>	No
<b>Conflict of interest</b>		Item 27: Describe sources of funding for the systematic review and other support (e.g., supply	Item 22. Give the source of funding and the role of the funders for the present study and, if applicable, for	Item 25. Sources of funding and other support (such as supply of drugs), role of funders



		of data); role of funders for the systematic review.	the original study on which the present article is based	
Private data computing company	Collecting / aggregating data by for-profit companies should be avoided. The financial links of such companies (shareholders) should be known.	Not mentioned	Not mentioned	Not mentioned
Centers and doctors identified	Centers and doctors recruiting patient should be known	Not mentioned	Item 5. Setting Describe the setting, locations, and relevant dates, including periods of recruitment, exposure, follow-up, and data collection	Item 4b. Settings and locations where the data were collected Item 10. Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions

Undeclared funding / conflict of interest	When investigating the funding of the study and conflicts of interest of each author <sup>1</sup> , no undeclared / indirect funding or conflict of interest should be found.	Research of funding is required but not research of conflict of interest	Research of funding is required but not research of conflict of interest	Research of funding is required but not research of conflict of interest
Potential conflict of interest	Study is not funded, and authors have not received fees from one or several pharmaceutical industries with a direct or indirect <sup>2</sup> conflict of interest with the results of the study	Not mentioned – PRISMA checklist does not mention conflict of interest of eligible studies.	Funding but not conflict of interest is considered.	Funding but not conflict of interest is considered.
<b>Clinical expertise</b>				
At least one of the authors is a clinical expert-in-the-field	For a viral respiratory infection such as Covid-19, at least 1 author is affiliated to an infectious disease, internal medicine or a pneumology unit.  Biomedical, public health specialists are not clinical experts as far as they do not care for patients.	Not mentioned	Not mentioned	Not mentioned

<p>Diagnosis is confirmed by a laboratory test</p>	<p>Patients without a laboratory test are excluded. Diagnosis should not rely on clinical or radiological evidence only.</p>	<p>Not mentioned</p>	<p>Item 6. Participants Give the eligibility criteria, and the sources and methods of case ascertainment and control selection. Item 7. Give diagnosis criteria, if applicable <i>No mention of a laboratory confirmation test.</i></p>	<p>Not mentioned</p>
<p>Detailed therapeutic protocol</p>	<p>A detailed therapeutic protocol is provided allowing a medical doctor expert-in-the-field to reproduce it, considering most common contraindications, precautions for use and monitoring.</p>	<p>Not mentioned</p>	<p>Not mentioned</p>	<p>Item 5. The interventions for each group are described with sufficient details to allow replication, including how and when they were actually administered.</p>

				<i>No mention of contraindications, precautions of use and monitoring</i>
Treatment is not toxic	Dose is usual, not in the overdose range and follows commonly-used doses with this drug. Drug is not used in patients with contraindications.	Not mentioned	Not mentioned	Not mentioned
A specific treatment effective on the microbe is not given to controls	Other drugs potentially effective on the microbe are known by the clinical expert. They should not be given to the untreated patients.	Not mentioned	Not mentioned	Not mentioned
Role of previous health status is ruled out	A difference in previous health status between treated and untreated should not interfere with outcome (typically a difference of age or mortality).  Combined Charlson score frequently used in this context.	Not mentioned	Item 7. Clearly define all outcomes, exposures, predictors, potential confounders, and effect modifiers.  Item 14. Give characteristics of study	Item 15. A table showing baseline demographic and clinical characteristics for each group

			<p>participants (e.g. demographic, clinical, social) and information on exposures and potential confounders</p> <p><i>Control for previous health status (at least age) not required</i></p>	
<p>Role of disease severity is ruled out</p>	<p>A difference in disease severity between treated and untreated should not interfere with outcome (typically a difference of vital parameters). NEWS score frequently used in this context.</p>	<p>Not mentioned</p>	<p>Item 7. Clearly define all outcomes, exposures, predictors, potential confounders, and effect modifiers.</p> <p>Item 14. Give characteristics of study participants (e.g. demographic, clinical, social) and information</p>	<p>Item 15. A table showing baseline demographic and clinical characteristics for each group</p>

			<p>on exposures and potential confounders</p> <p><i>Control for disease severity (at least the relevant vital parameters) not required</i></p>	
A 25% lower mortality should not be neglected	Authors should not conclude an absence of effect when risk of mortality is decreased of more than 25%	Not mentioned	Not mentioned	Not mentioned
An unexpected clinically relevant finding should not be neglected	Authors should report unexpected finding for instance a very different effect size in a subgroup	Item 16. Additional analyses. Describe methods of additional analyses (e.g., sensitivity or subgroup analyses, meta-regression), if done, indicating which were pre-specified.	Not mentioned	<p>Item 12b. Methods for additional analyses, such as subgroup analyses and adjusted analyses</p> <p>Item 18. Results of any other analyses performed, including</p>

				subgroup analyses and adjusted analyses, distinguishing pre-specified from exploratory
<b>Methodology</b>				
Design of the study (RCT/observational) is mentioned	Design of the study should be mentioned	Design of the study should be mentioned	Item 4. Study design. Present key elements of study design early in the paper	Item 1a. Identification as a randomized trial in the title
Big data studies are identified	If the data are analyzed by data scientists with electronic medical files without connection to the medical doctors who take care of patients, this should be mentioned	Not mentioned	Not mentioned	Not mentioned
Mono or multicentric design is known	This is naturally clarified when centers and doctors are known. If the study is an RCT with several recruiting centers, it should be identified as a megatrial	Not mentioned	Not mentioned	Not mentioned

	associated with specific risks of bias (Simpson's paradox). Effect should be reported for each center.			
Number of events and sample size of each group in each center are provided	This improves verifiability but is not sufficient <i>per se</i> to prevent other biases	Not mentioned	Not mentioned	For each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)  <i>No mention of the center</i>
In multicentric studies, adjusted effect is reported in each center	This prevents the Simpson's paradox	Simpson's paradox neglected	Simpson's paradox neglected	Simpson's paradox neglected

<sup>1</sup>For instance using governmental (<https://www.transparence.sante.gouv.fr>), or non-governmental transparency websites

(<https://projects.propublica.org/docdollars/>, <https://www.eurosfordocs.fr/>), <sup>2</sup>A direct conflict of interest is found when the drug tested is sold by the pharmaceutical company which funded the study and/or paid fees to one or several authors, an indirect conflict of interest is defined when the drug tested is in the same niche and in competition with a drug (or pharmaceutical product (i.e. a vaccine)) sold or developed by the pharmaceutical company which funded the study and/or paid fees to one or several authors.



**Supplementary Figure 1. One-study-removed meta-analysis of observational studies without potential conflict of interest and with detailed therapeutic protocol**

Study name	Statistics with study removed				Odds ratio (95% CI) with study removed						
	Point	Lower limit	Upper limit	p-Value	0.10	0.20	0.50	1.00	2.00	5.00	10.00
Alberici, Kidney International, 2020 - HCQ	0.61	0.52	0.70	0.000000			++				
Arshad, Int J Infect Dis, 2020 - HCQ +/- AZ	0.60	0.52	0.70	0.000000			++				
Ayerbe, Intern Emerg Med, 2020 - HCQ	0.61	0.52	0.71	0.000000			++				
Catteau, Int J Antimicrob Agents, 2020 - HCQ +/- AZ	0.57	0.48	0.69	0.000000			+				
Derwand, IJAA, 2020 - HCQ+AZ+Zinc	0.61	0.53	0.70	0.000000			++				
Di Castelnuovo, Eur J Intern Med, 2020 - HCQ	0.58	0.49	0.68	0.000000			++				
Guerin, Asian J Med Health, 2020 - HCQ+AZ	0.60	0.52	0.70	0.000000			++				
Lagier, Trav Med Infect Dis, 2020 - HCQ+AZ	0.60	0.52	0.70	0.000000			++				
Lauriola, Clin Translat Sci, 2020 - HCQ alone	0.60	0.52	0.69	0.000000			+				
Lauriola, Clin Translat Sci, 2020 - HCQ+AZ	0.65	0.58	0.74	0.000000			+				
Lecronier, Critical Care, 2020 - HCQ	0.61	0.53	0.70	0.000000			++				
Membrillo de Novales, Preprints, 2020 - HCQ	0.62	0.53	0.71	0.000000			++				
Mikami, J Gen Intern Med, 2020 - HCQ	0.59	0.50	0.69	0.000000			++				
Nachega, Am J Trop Med Hyg, 2020 - HCQ+AZ	0.64	0.56	0.73	0.000000			+				
Paccoud, Clin Infect Dis, 2020 - HCQ	0.60	0.52	0.70	0.000000			++				
Sulaiman, MedRxiv, 2020 - HCQ	0.57	0.49	0.67	0.000000			++				
Yu, Sci China Life Sci, 2020 - HCQ	0.61	0.53	0.71	0.000000			++				
	0.60	0.52	0.70	0.000000			++				

This analysis allows to exclude a summary significant effect linked to an aberrant study.

## References

1. Mehra MR, Desai SS, Ruschitzka F, Patel AN. RETRACTED: Hydroxychloroquine or chloroquine with or without a macrolide for treatment of COVID-19: a multinational registry analysis [published online ahead of print, 2020 May 22] [retracted in: Lancet. 2020 Jun 5;:null]. Lancet. 2020a;S0140-6736(20)31180-6. doi:10.1016/S0140-6736(20)31180-6
2. Million M, Gautret P, Colson P, et al. Clinical efficacy of chloroquine derivatives in COVID-19 infection: comparative meta-analysis between the big data and the real world. *New Microbes New Infect.* 2020;38:100709. Published 2020a Jun 6. doi:10.1016/j.nmni.2020.100709
3. Raoult D. Lancet gate: a matter of fact or a matter of concern. *New Microbes New Infect.* 2020;38:100758. doi:10.1016/j.nmni.2020.100758
4. Million M, Chaudet H, Raoult D. Hydroxychloroquine Failure: The End does not justify the Means [published online ahead of print, 2020 Aug 6]. *Clin Infect Dis.* 2020c;ciaa1117. doi:10.1093/cid/ciaa1117
5. Roussel Y, Million M, Chabrière E, Lagier JC, Raoult D. Be careful with Big Data: Re-analysis of Patient Characteristics and Outcomes of 11,721 Patients with COVID19 Hospitalized Across the United States [published online ahead of print, 2020 Oct 22]. *Clin Infect Dis.* 2020;ciaa1618. doi:10.1093/cid/ciaa1618
6. Million M, Roussel Y, Raoult D. Chloroquine and COVID-19: A western medical and scientific drift?. *Eur J Intern Med.* 2020;78:4-5. doi:10.1016/j.ejim.2020.06.020
7. Million M, Roussel Y, Lagier JC, Raoult D. Re: 'Effect of hydroxychloroquine with or without azithromycin on the mortality of COVID-19 patients' by Fiolet et al. *Clin Microbiol Infect.* 2021;27(1):132-133. doi:10.1016/j.cmi.2020.09.026

8. Raoult D. Rational for meta-analysis and randomized treatment: the COVID-19 example [published online ahead of print, 2020 Oct 21]. *Clin Microbiol Infect.* 2020a;S1198-743X(20)30643-1. doi:10.1016/j.cmi.2020.10.012
9. Fried MW, Crawford JM, Mospan AR, et al. Patient Characteristics and Outcomes of 11,721 Patients with COVID19 Hospitalized Across the United States [published online ahead of print, 2020 Aug 28]. *Clin Infect Dis.* 2020;ciaa1268. doi:10.1093/cid/ciaa1268
10. Wang Y, Zhang D, Du G, et al. Remdesivir in adults with severe COVID-19: a randomised, double-blind, placebo-controlled, multicentre trial [published correction appears in *Lancet.* 2020 May 30;395(10238):1694]. *Lancet.* 2020;395(10236):1569-1578. doi:10.1016/S0140-6736(20)31022-9
11. Boulware DR, Pullen MF, Bangdiwala AS, et al. A Randomized Trial of Hydroxychloroquine as Postexposure Prophylaxis for Covid-19. *N Engl J Med.* 2020;383(6):517-525. doi:10.1056/NEJMoa2016638
12. Skipper CP, Pastick KA, Engen NW, et al. Hydroxychloroquine in Nonhospitalized Adults With Early COVID-19: A Randomized Trial [published online ahead of print, 2020 Jul 16]. *Ann Intern Med.* 2020;M20-4207. doi:10.7326/M20-4207
13. Boschi C, Hoang VT, Giraud-Gatineau A, et al. Co-infections with SARS-CoV-2 and other respiratory viruses in Southeastern France: a matter of sampling time [published online ahead of print, 2020 Nov 24]. *J Med Virol.* 2020;10.1002/jmv.26692. doi:10.1002/jmv.26692
14. RECOVERY Collaborative Group, Horby P, Mafham M, et al. Effect of Hydroxychloroquine in Hospitalized Patients with Covid-19. *N Engl J Med.* 2020;383(21):2030-2040. doi:10.1056/NEJMoa2022926

15. Conan NJ Jr. The treatment of hepatic amebiasis with chloroquine. *Am J Med.* 1949;6(3):309-320. doi:10.1016/0002-9343(49)90167-9
16. Magagnoli J, Narendran S, Pereira F, et al. Outcomes of Hydroxychloroquine Usage in United States Veterans Hospitalized with COVID-19 [published online ahead of print, 2020 Jun 5]. *Med (N Y).* 2020;10.1016/j.medj.2020.06.001. doi:10.1016/j.medj.2020.06.001
17. McGrail DE, Dianna Edwards D. COVID-19 Case Series at UnityPoint Health St. Lukes Hospital in Cedar Rapids, IA medRxiv 2020.07.17.20156521; doi: <https://doi.org/10.1101/2020.07.17.20156521>
18. Alberici F, Delbarba E, Manenti C, et al. A report from the Brescia Renal COVID Task Force on the clinical characteristics and short-term outcome of hemodialysis patients with SARS-CoV-2 infection. *Kidney Int.* 2020;98(1):20-26. doi:10.1016/j.kint.2020.04.030
19. Mahévas M, Tran VT, Roumier M, et al. Clinical efficacy of hydroxychloroquine in patients with covid-19 pneumonia who require oxygen: observational comparative study using routine care data [published correction appears in *BMJ.* 2020 Jun 18;369:m2328]. *BMJ.* 2020;369:m1844. Published 2020 May 14. doi:10.1136/bmj.m1844
20. Geleris J, Sun Y, Platt J, et al. Observational Study of Hydroxychloroquine in Hospitalized Patients with Covid-19. *N Engl J Med.* 2020;382(25):2411-2418. doi:10.1056/NEJMoa2012410
21. Beigel JH, Tomashek KM, Dodd LE, et al. Remdesivir for the Treatment of Covid-19 - Final Report. *N Engl J Med.* 2020;383(19):1813-1826. doi:10.1056/NEJMoa2007764

22. Bernaola N, Mena R, Bernaola A, et al. Observational Study of the Efficiency of Treatments in Patients Hospitalized with Covid-19 in Madrid. medRxiv 2020.07.17.20155960; doi: <https://doi.org/10.1101/2020.07.17.20155960>
23. Ayerbe L, Risco-Risco C, Ayis S. The association of treatment with hydroxychloroquine and hospital mortality in COVID-19 patients [published online ahead of print, 2020 Sep 30]. Intern Emerg Med. 2020;1-6. doi:10.1007/s11739-020-02505-x
24. WHO Solidarity trial consortium, Hongchao Pan, Richard Peto, Quarraisha Abdool Karim, Marissa Alejandria, Ana Maria Henao-Restrepo, César Hernández García, Marie-Paule Kieny, Reza Malekzadeh, Srinivas Murthy, Marie-Pierre Preziosi, Srinath Reddy, Mirta Roses Periago, Vasee Sathiyamoorthy, John-Arne Røttingen, Soumya Swaminathan, as the members of the Writing Committee, assume responsibility for the content and integrity of this article. Repurposed antiviral drugs for COVID-19 – interim WHO SOLIDARITY trial results. medRxiv 2020.10.15.20209817; doi: <https://doi.org/10.1101/2020.10.15.20209817>
25. WHO Solidarity Trial Consortium, Pan H, Peto R, et al. Repurposed Antiviral Drugs for Covid-19 - Interim WHO Solidarity Trial Results [published online ahead of print, 2020 Dec 2]. N Engl J Med. 2020;NEJMoa2023184. doi:10.1056/NEJMoa2023184
26. El-Solh AA, Meduri UG, Lawson Y, Carter M, Mergenhagen KA. Clinical course and outcome of covid-19 acute respiratory distress syndrome: data from a national repository. medRxiv 2020.10.16.20214130; doi: <https://doi.org/10.1101/2020.10.16.20214130>
27. Flisiak R, Zarębska-Michaluk D, Berkan-Kawińska A, Tudrujek-Zdunek M, Rogalska M, Piekarska A, et al. Remdesivir-based therapy improved recovery of patients with

COVID-19 in the SARSTer multicentre, real-world study. medRxiv

2020.10.30.20215301; doi: <https://doi.org/10.1101/2020.10.30.20215301>

28. Spinner CD, Gottlieb RL, Criner GJ, et al. Effect of Remdesivir vs Standard Care on Clinical Status at 11 Days in Patients With Moderate COVID-19: A Randomized Clinical Trial. *JAMA*. 2020;324(11):1048-1057. doi:10.1001/jama.2020.16349
29. Tang W, Cao Z, Han M, et al. Hydroxychloroquine in patients with COVID-19: an open-label, randomized, controlled trial. medRxiv . 2020.04.10;20060558. doi: 10.1101/2020.04.10.20060558 (version 1).
30. Tang W, Cao Z, Han M, et al. Hydroxychloroquine in patients with mainly mild to moderate coronavirus disease 2019: open label, randomised controlled trial. *BMJ*. 2020;369:m1849. Published 2020 May 14. doi:10.1136/bmj.m1849
31. Magagnoli J, Narendran S, Pereira F, et al. Outcomes of Hydroxychloroquine Usage in United States Veterans Hospitalized with COVID-19 [published online ahead of print, 2020 Jun 5]. *Med (N Y)*. 2020;10.1016/j.medj.2020.06.001. doi:10.1016/j.medj.2020.06.001
32. Goldman JD, Lye DCB, Hui DS, et al. Remdesivir for 5 or 10 Days in Patients with Severe Covid-19 [published online ahead of print, 2020 May 27]. *N Engl J Med*. 2020;NEJMoa2015301. doi:10.1056/NEJMoa2015301
33. Cavalcanti AB, Zampieri FG, Rosa RG, et al. Hydroxychloroquine with or without Azithromycin in Mild-to-Moderate Covid-19 [published online ahead of print, 2020 Jul 23]. *N Engl J Med*. 2020;NEJMoa2019014. doi:10.1056/NEJMoa2019014
34. Mahévas M, Tran VT, Roumier M, et al. Clinical efficacy of hydroxychloroquine in patients with covid-19 pneumonia who require oxygen: observational comparative study using routine care data [published correction appears in *BMJ*. 2020 Jun

18;369:m2328]. *BMJ*. 2020;369:m1844. Published 2020 May 14. doi:

<https://doi.org/10.1136/bmj.m1844>

35. Clinical efficacy of hydroxychloroquine in patients with covid-19 pneumonia who require oxygen: observational comparative study using routine care data. *BMJ*. 2020;369:m2328. Published 2020 Jun 18. doi:10.1136/bmj.m2328
36. Lescure FX, Bouadma L, Nguyen D, et al. Clinical and virological data of the first cases of COVID-19 in Europe: a case series [published correction appears in *Lancet Infect Dis*. 2020 May 19;:] [published correction appears in *Lancet Infect Dis*. 2020 Jun;20(6):e116]. *Lancet Infect Dis*. 2020;20(6):697-706. doi:10.1016/S1473-3099(20)30200-0
37. Sallard E, Belhadi D, Lescure FX, Yazdanpanah Y, Peiffer-Smadja N. Clinical trial protocols of repurposed prophylaxis for COVID-19: A review [published online ahead of print, 2020 Oct 3]. *Med Mal Infect*. 2020;S0399-077X(20)30711-3. doi:10.1016/j.medmal.2020.09.013
38. Grein J, Ohmagari N, Shin D, et al. Compassionate Use of Remdesivir for Patients with Severe Covid-19. *N Engl J Med*. 2020;382(24):2327-2336. doi:10.1056/NEJMoa2007016
39. Dubert M, Visseaux B, Isernia V, et al. Case report study of the first five COVID-19 patients treated with remdesivir in France. *Int J Infect Dis*. 2020;98:290-293. doi:10.1016/j.ijid.2020.06.093
40. Lê MP, Peiffer-Smadja N, Guedj J, et al. Rationale of a loading dose initiation for hydroxychloroquine treatment in COVID-19 infection in the DisCoVeRy trial-authors' response. *J Antimicrob Chemother*. 2021;76(1):277-279. doi:10.1093/jac/dkaa415

41. Ip A, Berry DA, Hansen E, et al. Hydroxychloroquine and tocilizumab therapy in COVID-19 patients-An observational study. *PLoS One*. 2020;15(8):e0237693. Published 2020 Aug 13. doi:10.1371/journal.pone.0237693
42. Rivera DR, Peters S, Panagiotou OA, et al. Utilization of COVID-19 Treatments and Clinical Outcomes among Patients with Cancer: A COVID-19 and Cancer Consortium (CCC19) Cohort Study. *Cancer Discov*. 2020;10(10):1514-1527. doi:10.1158/2159-8290.CD-20-0941
43. Singh S, Khan A, Chowdhry M, Chatterjee A. Outcomes of Hydroxychloroquine Treatment Among Hospitalized COVID-19 Patients in the United States- Real-World Evidence From a Federated Electronic Medical Record Network. *medRxiv*. 2020.05.12;20099028; doi: 10.1101/2020.05.12.20099028
44. Annie FH, Sirbu C, Frazier KR, Broce M, Lucas BD Jr. Hydroxychloroquine in Hospitalized Patients with COVID-19: Real-World Experience Assessing Mortality. *Pharmacotherapy*. 2020;40(11):1072-1081. doi:10.1002/phar.2467
45. Sbidian E, Josse J, Lemaitre G, et al. Hydroxychloroquine with or without azithromycin and in-hospital mortality or discharge in patients hospitalized for COVID-19 infection: a cohort study of 4,642 in-patients in France. *medRxiv* 2020;06.16.20132597; doi: 10.1101/2020.06.16.20132597
46. Kelly M, O'Connor R, Townsend L, et al. Clinical outcomes and adverse events in patients hospitalised with COVID-19, treated with off-label hydroxychloroquine and azithromycin [published online ahead of print, 2020 Jul 20]. *Br J Clin Pharmacol*. 2020;10.1111/bcp.14482.
47. Peters EJG, Collard D, Sander van Assen S, et al. Outcomes of Persons With COVID-19 in Hospitals With and Without Standard Treatment With (Hydroxy)chloroquine. *medRxiv* 2020.08.14.20173369; doi: <https://doi.org/10.1101/2020.08.14.20173369>



48. Alamdari NM, Afaghi S, Rahimi FS, et al. Mortality Risk Factors among Hospitalized COVID-19 Patients in a Major Referral Center in Iran. *Tohoku J Exp Med.* 2020;252(1):73-84. doi:10.1620/tjem.252.73
49. Bhandari S, Singh A, Sharma R, et al. Characteristics, Treatment Outcomes and Role of Hydroxychloroquine among 522 COVID-19 hospitalized patients in Jaipur City: An Epidemio-Clinical Study. *J Assoc Physicians India.* 2020;68(6):13-19.
50. Brouqui P, Amrane S, Million M, et al. Asymptomatic hypoxia in COVID-19 is associated with poor outcome [published online ahead of print, 2020 Oct 31]. *Int J Infect Dis.* 2020;102:233-238. doi:10.1016/j.ijid.2020.10.067
51. Çalik BaŞaran N, Uyarođlu OA, Telli Dizman G, et al. Outcome of Non-Critical COVID-19 Patients with Early Hospitalization and Early Antiviral Treatment Outside the ICU [published online ahead of print, 2020 Jul 28]. *Turk J Med Sci.* 2020;10.3906/sag-2006-173. doi:10.3906/sag-2006-173
52. Derwand R, Scholz M, Zelenko V. COVID-19 outpatients: early risk-stratified treatment with zinc plus low-dose hydroxychloroquine and azithromycin: a retrospective case series study. *Int J Antimicrob Agents.* 2020;56(6):106214. doi:10.1016/j.ijantimicag.2020.106214
53. Heberto AB, Carlos PCJ, Antonio CRJ, et al. Implications of myocardial injury in Mexican hospitalized patients with coronavirus disease 2019 (COVID-19). *Int J Cardiol Heart Vasc.* 2020;30:100638. doi:10.1016/j.ijcha.2020.100638
54. Guérin V, Lévy P, Thomas, J, Lardenois T, et al. Azithromycin and Hydroxychloroquine Accelerate Recovery of Outpatients with Mild/Moderate COVID-19. *Asian J Med Health* 2020;18:45-55. doi: 10.9734/ajmah/2020/v18i730224

55. Pinato DJ, Zambelli A, Aguilar-Company J, et al. Clinical portrait of the SARS-CoV-2 epidemic in European cancer patients [published online ahead of print, 2020 Jul 31]. *Cancer Discov.* 2020;CD-20-0773. doi:10.1158/2159-8290.CD-20-0773
56. Roomi S, Ullah W, Ahmed F, et al. Efficacy of Hydroxychloroquine and Tocilizumab in Patients With COVID-19: Single-Center Retrospective Chart Review. *J Med Internet Res.* 2020;22(9):e21758. Published 2020 Sep 1. doi:10.2196/21758
57. Serrano Domingo JJ, Corral de la Fuente E, Martin Huertas R, Vida Navas EM, Soto Castillo JJ, Sanz Gomez L, et al. COVID19 in cancer patients: Risk factors for the development of severe clinical events (SCE). 2020;31 suppl 4, S1019 (1744P) doi : 10.1016/j.annonc.2020.08.1808
58. Soto-Becerra P, Culquichicon C, Hurtado-Roca Y, Araujo-Castillo RV. Real-World Effectiveness of hydroxychloroquine, azithromycin, and ivermectin among hospitalized COVID-19 patients: Results of a target trial emulation using observational data from a nationwide Healthcare System in Peru. medRxiv 2020.10.06.20208066; doi: <https://doi.org/10.1101/2020.10.06.20208066>
59. Lagier JC, Million M, Gautret P, et al. Outcomes of 3,737 COVID-19 patients treated with hydroxychloroquine/azithromycin and other regimens in Marseille, France: A retrospective analysis. *Travel Med Infect Dis.* 2020;36:101791. doi:10.1016/j.tmaid.2020.101791
60. Leger T, Jacquier A, Barral PA, et al. Low-dose chest CT for diagnosing and assessing the extent of lung involvement of SARS-CoV-2 pneumonia using a semi quantitative score. *PLoS One.* 2020;15(11):e0241407. Published 2020 Nov 3. doi:10.1371/journal.pone.0241407
61. Sulaiman T, Mohana A, Alawdah L, et al. The Effect of Early Hydroxychloroquine-based Therapy in COVID-19 Patients in Ambulatory Care Settings: A Nationwide

Prospective Cohort Study. medRxiv 2020.09.09.20184143; doi:

<https://doi.org/10.1101/2020.09.09.20184143>

62. Synolaki E, Papadopoulos V, Divolis G, et al. Activin/Follistatin-axis deregulation is independently associated with COVID-19 in-hospital mortality. medRxiv 2020.09.05.20184655; doi: <https://doi.org/10.1101/2020.09.05.20184655>